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Chavent, G.: Analyse fonctionnelle et identification de coefficients répartis dans les équations aux dérivées partielles, Thése d'Etat, Facultè des Sciences de Paris, 1971. Richter, G.R.: An inverse problem for the steady state diffusion equation, SIAM J.

Richter, G.R.: Numerical identification of spatially varying diffusion coefficient, Mathematics of Computation, 36(1981), 375-386.

Emsellem, Y and G. de Marsily. An automatic solution for the inverse problem, Water

Resour. Res., 7(1971), 1264-1283. Chicone, C. and J. Gerlach: A note on the identifiability of distributed parameters in

elliptic equations, SIAM J. Math. Anal., 18(1987), 1378-1384. Sagar, B., Yakowitz, S. and L. Duckstein: A direct method for the identification of the parameters of dynamic nonhomogeneous aquifers, Water Resour. Res., 11(1975), 563-

570.

Parravicini, G., Giudici, M., Morossi, G. and G. Ponzini: Minimal assignment of phenomenological coefficients and uniqueness for an inverse problem, Preprint IFUM 480/FT, Dipartimento di Fisica, Università di Milano, Milan, Italy, (1994).

Giudioi, M., Morossi, G., Parravicini, G. and G. Ponzini: A new method for the identification of distributed transmissivities, Preprint IFUM 481/FT, Dipartimento di Fisica, Università di Milano, Milan, Italy, (1994).

Samarskij, A. and V. Andreev: Méthodes aux différences pour equation elliptiques, Mir,

Moskow 1978.

Carrera, J. and S.P. Neuman: Estimation of aquifer parameters under transient and steady-state conditions: 3. application to synthetic and field data, Water Resour. Res.,

22(1986), 228-242.
Scarascia, S. and G. Ponzini: An approximate solution for the inverse problem in budrantics 1/Bneroia Flettrica 49(1972) 518-531.

hydraulics, L'Energia Elettrica, 49(1972), \$18-531.

Carrera, J. and S.P. Neuman: Estimation of Aquifer parameters under transient and steady-state conditions: 1, maximum likelihood method incorporating prior information, Water Resour. Res., 22(1986), 199-210.

Carrera, J. and S.P. Neuman: Estimation of aquifer parameters under transient and steady-state conditions: 2. uniqueness, stability and solution algorithms, Water Resour. Res. 27(1986) 211-227

Emsellen, Y. and G. de Marsily. Reply to "Comments on 'An authomatic solution for the inverse problem' by Y. Emsellem and G. de Marsily" by D. Kleinecke, Water Resour. Res, 8(1972), 1130-1131.

VERIFICATION OF ACTIVE AND PASSIVE GROUND-WATER CONTAMINATION REMEDIATION EFFORTS

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ABSTRACT

The verification of ground-water contamination remediation efforts requires thorough documentation of subsurface conditions before, during and after cleanup efforts have ceased. The documentation include include proof of: reduction of risk to human or environmental health, achievement of regulatory cleanup concentration goals in soil, gas or liquid media, or verification of continued approach to background environmental quality conditions. Meeting any one or all of these requirements calls for a comprehensive approach to the design and operation of remediation efforts with an emphasis on the monitoring of environmental conditions. These tasks are most challenging for in-situ remediation efforts which employ active (i.e., pumping or vacuum application) rather than passive (i.e., natural water and vapor gradient)

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1. INTRODUCTION

The practice of site characterization for potential organic contaminants has evolved slowly in the past decade. Early guidelines [1,2,3], for minimal ground-water contamination detection nonitoring (i.e., monitoring wells upgradient and downgradient) have been applied to many sites of potential concern from detection through remedial action selection phases.

physicochemical characteristics of contaminant mixtures or the complexity substantially more comprehensive approaches. For organic contaminant remediation principally by in-situ biological methods for volatile organic organic components of fuel and solvent mixtures have generated a flurry detection and assessment (i.e., determination of the nature and extent of of modified, monitoring well-based site characterization approaches [4]. This minimal approach has often been applied regardless of the contamination) efforts, wells alone have been found to be inadequate Recognition of the value of subsurface soil vapor surveys for volatile of the hydrogeologic setting. For soluble inorganic constituents, this approach may be adequate for detection. Assessment efforts require However, these approaches to site characterization and monitoring monitoring tools. This paper focuses on the monitoring needs for network design suffer from the failure to identify the total mass of compounds present in hydrocarbon fuels and organic solvents. contaminant in the subsurface for three main reasons. First, although volatile organic compounds (VOC's) are mobile in ground-water and frequently early indicators of plume movement [5], their detection in vapor or well samples and apparent aqueeus concentration distribution does not identify the total mass distribution of organic contaminant [6]. Secondly, efforts to correlate observed soil vapor or ground water VOC concentrations with those in subsurface solid cores have often been unsuccessful. This is because current bulk jar collection/refrigeration at 4°C guidelines for solid core samples for VOC analyses lead to gross negative errors [7]. Thirdly, "snapshots" (i.e, one-time surveys) of background and disturbed ground-water chemistry conditions have been interpreted as "constant" ignoring temporal variability in subsurface geochemistry.

The result of the slow improvement in site-characterization and monitoring practices has often been the very low probability detection of the source of mobile organic contaminants. This outcome may be followed by the misapplication of risk-assessment and remediation models.

Nonetheless, there exist good reasons for a more optimistic view for the future reliability of site characterization and monitoring efforts.

2. ACTIVE AND PASSIVE REMEDIATION APPROACHES

the volume of the treatment zone, these monitoring data provide the basis process stream and in-situ environmental conditions for concentrations of hydraulic (ground-water) or pneumatic (vapor) control within the zone of of a suite of physical, chemical, and microbiological processes to destroy or subsurface ground-water or vapor flow. Also they rely on the application calls for careful design of an active remediation-based monitoring system. contaminants. The reliability of these data is critical to the verification of Active in-situ remediation efforts generally involve the control of parent compounds and transformation products. Linked to net flow and treatment. This facilitates contaminant removal or transformation but treatment designs where extracted fluids are returned to the subsurface Such levels of control have most often been achieved in "closed-loop" treatment zone. In these instances it is necessary to monitor both the constituents. The most effective active remediation schemes sustain transform contaminants to less harmful or less mobile chemical for estimates of the net removal/transformation of the original cleanup performances.

Passive remediation efforts rely on intrinsic biological and/or chemical processes to mediate the destruction or transformation of contaminants. Though they may take more time to achieve acceptable levels of contaminant removal than active methods, the existing monitoring design from detective or assessment phases of the project may need only slight modification as to sampling location, frequency and selection of monitoring parameters. This approach may significantly reduce the cost of remediation.

The shortcomings of previous contaminant detection and assessment monitoring efforts have been recognized. New guidelines and recommendations on network design and operations will lead to more comprehensive, cost-effective site characterization [7,9] in general. Also, excellent reviews of characterization and long term monitoring needs and approaches in support of in-situ remediation efforts should guide us in this regard [10,11]. Site characterization efforts provide a basis for long term monitoring design and actually continue throughout the life of a remediation project.

Active and passive in-situ bioremediation approaches have been applied frequently to subsurface cleanups of organic contaminants (e.g., fuels, solvents, pesticides, etc.). The monitoring measures for verification of bioremediation performances have been identified as: the documented decline of contaminant concentrations, identification of favorable conditions (e.g. substrate, nutrient, pH, electron-acceptors) for microbial activity, demonstration of an active microbial population capable of transforming the major contaminants, and the identification of intermediate break-down or end-products in the subsurface. The supply of suitable electron acceptors (e.g. O2, NO3-, FeIII, SO4=, etc.) may be the crucial element in successful in-situ remediation efforts.

While these measures are necessary, they are not sufficient to establish the remedial effectiveness or performance of in-situ methods. The minimal measures noted above must be integrated into a mass-balance for contaminants and transformation products. A number of inorganic and organic indicators of subsurface transformation can be used to permit the approach to mass balance for specific organic contaminants. Table 1 shows various example monitoring indicators appropriate for solvent and fuel contamination situations where transformations occur under known limits of oxidation-reduction conditions.

Organic Specific	aromatic acids	Trichloro- ethylene, Trichloro- ethane Dichloro- ethylenes Vinyl chloride C2H4-ethylene C3Hs-ethane
Organic General	low- molecular wt. organic acids	low- Trichloro- molecular wt. ethylene, organic acids Trichloro- CH4-methane ethane Dichloro- ethylenes Vinyl chlo C2H4-ethyl
Inorganic	CO ₂ O ₂ NO ₃ - Fell	CO, N.I.3 Fell
Contaminant Mixture	Gasoline GO ₂ (Benzene, O ₂ Toluene, NO ₃ Xylene, NO ₂ Alkylbenzenes Fell	Tetrachloro- ethylene Trichloro- ethylene
General Conditions	aerobic (oxic)	anaerobic (anoxic)

TABLE 1 General Monitoring Indicators for Organic Contaminant Mixtures

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It should be noted that subsurface redox conditions are not in chemical equilibrium and that transitional environments exist where intermediate transformation product stability may be significant [8].

The use of these indicators along with monitoring the concentration of the original compounds provides a more comprehensive approach to verifying remediation performance. There are relatively few examples of the mass balance approach, since there may be multiple pathways for field microbial transformation and the reaction products may be unknown. The pathways for microbial transformation are being delimited by a combination of field and laboratory experiments. Also the suite of reaction products are being determined by advanced analytical methods which will support the mass balance approach.

ADVANCED SITE CHARACTERIZATION AND MONITORING

How do we proceed to estimate the potential for subsurface intrinsic bioremediation success and track its performance into the future? Clearly, we should seek to design technically-defensible characterization and monitoring networks which will provide reasonable estimates of the inplace contaminant distribution over time. A dynamic, ongoing site characterization effort therefore includes objectives to:

- identify the spatial distribution of contaminants, particularly their relative fractionation in subsurface solids, water, and vapor, along potential exposure pathways recognizing that the mass of contaminants frequently resides in the solids;
 - 2) determine the corresponding spatial distribution of total organic matter since overall microbial activity and disruptions in subsurface geochemical conditions (and bioremediation indicators) are due to the total mass of reactive organic carbon;
- 3) estimate the temporal stability of hydrogeologic and geochemical conditions which may favor microbial transformations in background, source and downgradient zones during the first year of characterization and monitoring;
 - 4) derive initial estimates of net microbial transformations of contaminant-related organic matter over time which may be built into the long-term monitoring network design.

The first three objectives establish the environment of major contamination and the conditions under which bioremediation may occur. The latter two objectives are vitally important since the evaluation of the progress of intrinsic bioremediation processes depends on distinguishing

Verification of Active and Passive Ground-Water Contamination

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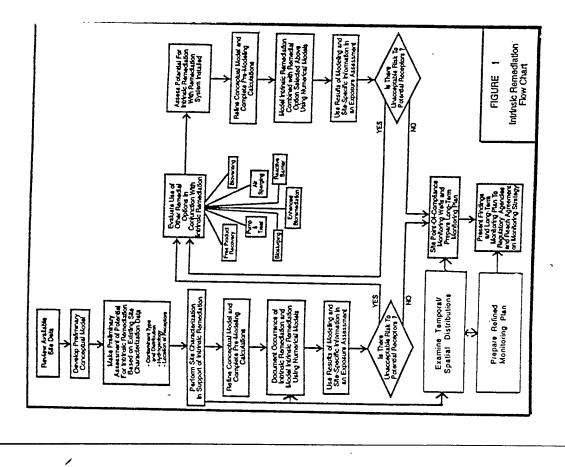
microbial transformations. This approach has been suggested emphatically compound "losses" due to dilution, sorption and chemical reactions from by Wilson [10] and was recently developed into a technical U.S. Air Force (ÚSAF) protocol by Wiedemeier, et al. [11].

the original work. The USAF Protocol [11] has as its goals the collection of The general approach is shown in Figure 1 which has been modified from intrinsic remediation for dissolved fuel contamination in ground water. The latter reference focuses directly on the implementation of data necessary to support:

- 1) Documented loss of contaminants of the field scale, 2) The use of chemical analytical data in mass balance calculations,
- Laboratory microcosm studies using aquifer samples collected from the site. 3

decision tool depends on the complexity of the actual hydrogeologic setting collection and analysis. The major categories of necessary data are listed in aquifer property, hydrogeologic or geochemical data needed to formulate a time should be sufficient to successfully implement intrinsic remediation refined and used to determine the suitability of intrinsic remediation as a These data, if collected in three dimensions for an extended period support the development of a site-specific conceptual model. This model characterization database. The USAF Protocol is quite comprehensive in available prior to in-depth site characterization are more likely to contain [12]. The data collected in the initial site characterization effort (Figure 1) geochemical data for a site. The conceptual model, in turn, can be tested, risk-management strategy. The validity of the conceptual model as a contaminant-related information rather than the three-dimensional conceptual model. A recognition of the variability inherent in these Table 2 from the USAF Protocol [11]. Detective monitoring datasets is a three dimensional representation of the ground water flow and and contaminant distributions relative to the completeness of the identifying important parameters, inputs and procedures for data parameter distributions is critical to site-characterization efforts. transport fields based on geologic, hydrologic, climatologic, and

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FRACTIONATION AND SPATIAL EXTENT OF CONTAMINATION

- Extent and type of soil and ground water contamination
 - (i.e., areas containing free- or residual-phase product) 2. Location and extent of contaminant source area(s)

1

The potential for a continuing source due to leaking tanks or pipelines

HYDROGEOLOGIC AND GEOCHEMICAL FRAMEWORK

- 4. Ground water geochemical parameter distributions (Table 3)
 - Regional hydrogeology including:
- Drinking water aquifers and
- Regional confining units.
- 6. Local and site-specific hydrogeology, including:
 - Local drinking water aquifers;
- Location of industrial, agricultural, and domestic water
- Patterns of aquifer use;
 - Lithology;
- Site stratigraphy, including identification of transmissive
 - Grain-size distribution (sand vs. silt vs. clay); and nontransmissive units;
- Aquifer hydraulic conductivity determination and estimates from grain-size distributions;
 - Ground water hydraulic information;
- Preferential flow paths;
- Location and type of surface water bodies; and
- Areas of local ground water recharge and discharge.
- Definition of potential exposure pathways and receptors.

TABLE 2

Site Specific Parameters to be Determined during Site Characterization (modified from Reference 11)

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SAMPLING IN SPACE 3.1

Verification of Active and Passive Ground-Water Contamination

contaminants, aquifer properties, and geochemical constituents (Table 3) derived from spatial averages of data points must include approximately 30 or more data points [13,14,15]. Indeed, this minimum dataset size strictly The initial site characterization phase should be designed to provide spatially dense coverage of critical data over volumes corresponding to ten donors (e.g., organic carbon, Fe²⁺, S⁼, NH3, etc.) and electron acceptors (e.g. with regard to how spatially averaged masses of contaminants, electron applies to points in a plane. Two major decisions which must be made to one-hundred year travel times along ground water flow paths. The "volume-averaged" values of the contaminants, hydrogeologic and geochemical parameters within zones along the flow path(s) should be Specifically, this means that the datasets for derived mass loadings of derived from large enough datasets to permit estimation of statistical properties (e.g., mean, median, correlation distance, variance, etc.). O2, NO3, NO2, Fe and Mn oxides, SO4, etc.) are to be estimated.

grain size, laboratory estimates of hydraulic conductivity, etc.) the answer is constituents particularly VOC's which are sparingly water soluble, the bulk of the contaminant mass may in fact reside in the solids though both solids The first question deals with identification of the media in which the bulk of the constitutent's mass resides. For aquifer properties (e.g., simple. In this case, the solids are clearly the media of interest. For and water samples must be collected carefully.

decommissioned Wurtsmith AFB near Oscoda, Ml. In this case the bulk of technologies and/or multilevel sampling devices present very useful tools order to approach this level of depth detail in sampling, the use of "push" sampling of water should be considered in many VOC investigations. In watertable. In this situation, averaging data points over depths of > 0.5m site. Continuous coring of subsurface soids and close interval (i.e., < 1 m) associated concentration differences. This type of situation is typified by could easily lead to order of magnitude errors in estimated masses for a fringe/water table interface typically exhibits order of magnitude solidthe contaminant mass along the axis of a dissolved BTEX plume with the BTEX data shown in Figure 2 for a fire training area at the recently The second question pertains to the depth interval over which concentrations less than $1000 \, \mu g/L$ resides in aquifer solids below the "planar" data points may be averaged. With fuel-related aromatic contaminants the depth interval above and below the capillary for site characterization.

CONTAMINATION	APPARENT/	CONTAMINANT	INORGANIC CONSTITUENTS	INTRINSIC
AREA	GEOCHEMICAL	MIXTURE	İ	CONSTITUENTS
	REDOX ZONE			
SOURCE	REDUCING ANOXIC	FUELS CHLORINATED SOLVENTS	O ₂ , CO ₂ , H ₂ S; pH Fe ²⁺ , HS ⁻ /S ⁼ , NO ₂ ⁻ , NH ₃ , ALKALINITY	ORGANIC CARBONS, CH ₄ ORGANIC ACIDS PHENOLS AS ABOVE AND: CHLORINATED METABOLITES ETHYLENE, ETHANE
DOWNGRADIENT	TRANSITIONAL/ SUBOXIC	FUELS CHLORINATED SOLVENTS	O ₂ , CO ₂ , H ₂ S; pH, Fe ²⁺ ALKALINITY, NO ₂ , NO ₃ , NH ₃ , HS ⁻ /S ⁼	ORGANIC CARBON, CH ₄ ORGANIC ACIDS PHENOLS AS ABOVE AND: CHLORINATED METABOLITES ETHYLENE, ETHANE
UPGRADIENT/FAR- FIELD DOWNGRADIENT	OXIC	FUELS CHLORINATED SOLVENTS	O ₂ , CO ₂ , H ₂ S ALKALINITY, Fe ²⁺ , NO ₃ , NO ₂ , NH ₃	ORGANIC CARBON, CH4 ORGANIC ACIDS PHENOLS AS ABOVE AND: CHLORINATED METABOLITES ETHYLENE, ETHANE

TABLE 3 Target Constituents for Site Characterization in Support of Intrinsic Bioremediation

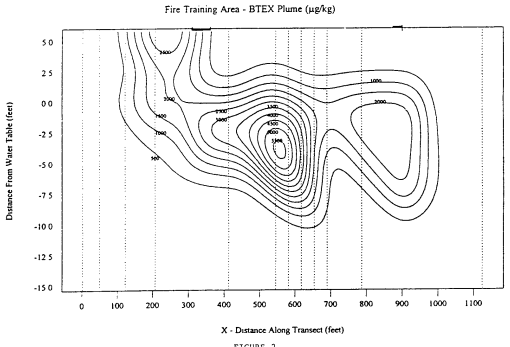


FIGURE 2

The approach to site characterization for chlorinated hydrocarbons is significantly more difficult. There are very few models of site characterization for these contaminants which have estimated mass loadings in specific media. Many of the previously referenced methods may work satisfactorily. However, free-phase detection, assessment and quantitation may be expected to be more a matter of luck and exhaustive sampling rather than intuition based on experience.

3.3 SAMPLING OVER TIME

VOC compounds (e.g., aromatic hydrocarbons, chlorinated solvents) are among the target contaminants which have been considered as constituents of concern in remedial investigations. Their aqueous solubility and demonstrated association with aquifer solids requires sampling of these media during the site characterization phase. This suggestion also applies to organic metabolites of complex organic mixtures (e.g., ethylene, vinyl chloride, aromatic acids, phenols). Aqueous plumes which develop subsequent to the release of these organic mixtures and byproduct compounds have received the most attention in the past. The fact that the mass of these contaminants frequently resides in the solids strongly suggests that the solids should receive the most attention in the initial site characterization effort. This should also be the case for the physical, geochemical, and microbial determinations.

Initially, conventional nested monitoring wells with screened lengths of 1 meter or more will be useful for estimating the spatial extent of the dissolved plume, for delineating apparent geochemical zones and to provide water level and aquifer property (e.g., slug and pump test derived hydraulic conductivity estimates). Semi-annual to annual sampling of wells, particularly multilevels appropriately designed and completed, should be quite useful over the course of the long term monitoring program. In this vein, their use should track the downgradient progress of risk-associated target compounds and permit testing predictions of intrinsic bioremediation effects on risk reduction.

However, proof of the effects of the net removal of specific solid-associated contaminants due to intrinsic bioremediation will depend on solid sampling and analysis at annual intervals or greater. This is because solid-associated concentration may be expected to change slowly. Unless biotransformation can be shown to be a major loss mechanism for contaminants mainly in solids over extended periods of time it will remain an area of research rather than practice.

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Since very few contamination situations have been monitored intensively for periods exceeding several years, it is difficult to define specific sampling frequencies for the range of hydrogeologic and contaminant combinations which may be encountered. Suffice it to say that the adoption and future refinement of technically defensible protocols which have been developed recently will improve intrinsic remediation approaches to risk management in subsurface contamination situations.

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REFERENCES

- Scalf, M.R., J.F. McNabb, W.J. Dunlop, R.L. Cosby, and J.S. Fryberger: Manual of Groundwater Sampling Procedures, National Water Well Association, (1981).
- Barcelona, M.J., J.P. Gibb, J.A. Helfrich and E.E. Garske: Practical Guide for Ground-Water Sampling. Illinois State Water Survey, SWS Contract Report 374, Ada, OK: U.S. Environmental Protection Agency (1985a).
- 3. U.S. Environmental Protection Agency, Office of Waste Programs Enforcement, Office of Solid Waste and Emergency Response, RCRA Technical Enforcement Guidance Document, OSWER-9950.1, U.S. Government Printing Office, Washington, D.C., (1986), 208 pp. Appendices.
- Eklund, B.: Detection of Hydrocarbons in Ground Water by Analysis of Shallow Soil Gas/Vapor, API Publication No. 4394, Washington, D.C., (1985).
- 5. Plumb, R.H.: A comparison of ground-water monitoring data from CERCLA and RCRA sites, Ground Wat. Mon. Rev., 7 (1987), 94-100.

Constitution of the second of

- 6. Robbins, G.A.: Influence of using purged and partially penetrating wells on contaminant detection, mapping and modeling, Ground Wat., 27 (1989), 155-162.
- 7. U.S. Environmental Protection Agency, Office of Solid Waste, RCRA Ground Water Monitoring: Draft Technical Guidance Document, U.S. Environmental Protection Agency, Washington, D.C., (1992a) EPA /530-R-93-001
- Barcelona, M.J., T. R. Holm, M.R. Scheck and G.E. George: Spatial and temporal gradients in aquifer oxidation-reduction conditions, Wat. Res. Res., 25 (1989), 998-1003.
- U.S. Environmental Protection Agency, Proceedings of the Ground Water Sampling Workshop, Dallas, TX, December 8-10, 1993. U.S. EPA-R.S. Kerr Laboratory, Ada, OK, EPA Office of Solid Waste, Washington, D.C., (1994).
- Wilson, J.T. 1993: Testing Bioremediation in the Field, p. 160-184 in In-Situ Bioremediation - When Does It Work?, Committee on In-Situ Bioremediation, Water Science and Technology Board National Research Council, National Academy Press, Washington, D.C. 207 pp.
- 11. Personal communication. T.H. Wiedemeier T.H. Wiedemeier, D.C. Downey, J.T. Wilson, D.H. Kampbell, R.N. Miller, J.E. Hansen, Draft Technical Protocol for Implementing the Intrinsic Remediation (Natural Attenuation) with Long-Term Monitoring Option for Dissolved-Phase Fuel Contamination in Ground Water. Air Force Center for Environmental Excellence, Brooks AFB, San Antonio, TX, March 1994.
- 12. National Research Council, In-Situ Bioremediation -When Does It Work?, National Academy Press, Washington, D.C. (1993), 207 pp.
- 13. Journel, A.G.: Geostatistics: models and tools for the earth sciences, Math. Geol., 18 (1986), 119-140.
- 14. Hoeksema, R.J. and P.K. Kitanidis: Analysis of the spatial structure of properties of selected aquifers, Wat. Res., Res., 21 (1985), 563-572.

Verification of Active and Passive Ground-Water Contamination

 Gilbert, R.O. and J.C Simpson.: Kriging for estimating spatial patterns of contaminants: potential and problems," Environ. Monit. and Assess., 5 (1985), 113-135.

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ADVANCED METHODS FOR GROUNDWATER POLLUTION CONTROL

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